# UTILITY PATENT APPLICATION TRANSMITTAL (Large Entity)

(Only for new nonprovisional applications under 37 CFR 1.53(b))

Docket No. BSI-469US

Total Pages in this Submission 49

### TO THE ASSISTANT COMMISSIONER FOR PATENTS

Box Patent Application Washington, D.C. 20231

Transmitted herewith for filing under 35 U.S.C. 111(a) and 37 C.F.R. 1.53(b) is a new utility patent application for an invention entitled:

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## UTILITY PATENT APPLICATION TRANSMITTAL (Large Entity)

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### **Application Elements (Continued)** ☑ Drawing(s) (when necessary as prescribed by 35 USC 113) a. 🔲 Formal Number of Sheets b. 🗵 Informal Number of Sheets Oath or Declaration a. 🔀 Newly executed (original or copy) ☐ Unexecuted b. Copy from a prior application (37 CFR 1.63(d)) (for continuation/divisional application only) c. With Power of Attorney ☐ Without Power of Attorney d. DELETION OF INVENTOR(S) Signed statement attached deleting inventor(s) named in the prior application, see 37 C.F.R. 1.63(d)(2) and 1.33(b). ☐ Incorporation By Reference (usable if Box 4b is checked) The entire disclosure of the prior application, from which a copy of the oath or declaration is supplied under Box 4b, is considered as being part of the disclosure of the accompanying application and is hereby incorporated by reference therein. 6. ☐ Computer Program in Microfiche (Appendix) 7 ☐ Nucleotide and/or Amino Acid Sequence Submission (if applicable, all must be included) a. Paper Copy b. Computer Readable Copy (identical to computer copy) с. 🔲 Statement Verifying Identical Paper and Computer Readable Copy **Accompanying Application Parts** Assignment Papers (cover sheet & document(s)) ☐ 37 CFR 3.73(B) Statement (when there is an assignee) ☐ English Translation Document (if applicable) 11. Information Disclosure Statement/PTO-1449 Copies of IDS Citations 12. Preliminary Amendment 13. Acknowledgment postcard 14. Certificate of Mailing First Class 🗵 Express Mail (Specify Label No.): EL684159501US

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	Accompanying Application Parts (Continued)												
15.		Certified Copy of Priority Document(s) (if foreign priority is claimed)											
16.		Additional Enclosures (please identify below):											
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<ul> <li>A check in the amount of \$1,196.00 to cover the filing fee is enclosed.</li> <li>☑ The Commissioner is hereby authorized to charge and credit Deposit Account No. 18-0350 as described below. A duplicate copy of this sheet is enclosed.</li> <li>☐ Charge the amount of as filing fee.</li> <li>☑ Credit any overpayment.</li> <li>☑ Charge any additional filing fees required under 37 C.F.R. 1.16 and 1.17.</li> <li>☐ Charge the issue fee set in 37 C.F.R. 1.18 at the mailing of the Notice of Allowance, pursuant to 37 C.F.R. 1.311(b).</li> </ul>													
Signature  Christopher R. Lewis, Reg. No. 36,201  Attorney for Applicant							01						
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### COMBINATION SELF-EXPANDABLE, BALLOON-EXPANDABLE ENDOLUMINAL DEVICE

#### TECHNICAL FIELD

This invention relates generally to endoluminal devices such as vena cava filters, stents, grafts, and/or prostheses and, more specifically, to endoluminal devices that have combined self-expanding and balloon-expandable properties.

#### BACKGROUND OF THE INVENTION

A stent is an elongated device used to support an intraluminal wall. In the case of a stenosis, a stent provides an unobstructed conduit for blood in the area of the stenosis. Such a stent may also have a prosthetic graft layer of fabric or covering lining the inside or outside thereof, such a covered stent being commonly referred to in the art as an intraluminal prosthesis, an endoluminal or endovascular graft (EVG), or a stent-graft.

A prosthesis may be used, for example, to treat a vascular aneurysm by removing the pressure on a weakened part of an artery so as to reduce the risk of rupture. Typically, a prosthesis is implanted in a blood vessel at the site of a stenosis or aneurysm endoluminally, i.e. by so-called "minimally invasive techniques" in which the prosthesis, restrained in a radially compressed configuration by a sheath or catheter, is delivered by a deployment system or "introducer" to the site where it is required. The introducer may enter the body through the patient's skin, or by a "cut down" technique in which the entry blood vessel is exposed by minor surgical means. When the introducer has been threaded into the body lumen to the prosthesis deployment location, the introducer is manipulated to cause the prosthesis to be ejected from the surrounding sheath or catheter in which it is restrained (or alternatively the surrounding sheath or catheter is retracted from the prosthesis), whereupon the prosthesis expands to a predetermined diameter at the deployment location, and the

introducer is withdrawn. Stent expansion may be effected by spring elasticity, balloon expansion, or by the self-expansion of a thermally or stress-induced return of a memory material to a pre-conditioned expanded configuration.

A vena cava filter is an implantable filter, typically implanted in a patient's inferior vena cava using minimally invasive techniques, for reducing the risk of arterial plaques dislodged during surgery or clots formed in the bloodstream from becoming lodged in and partially or completely blocking vessels supplying blood and oxygen to critical organs, such as the heart, lungs and brain. Such filters enable dangerous blood clots to be removed from circulation and held in a safe location until they can be dissolved by medication or extracted, again using minimally invasive techniques. Conventional implantable blood filters employing a variety of geometries are known, but many are generally basket or cone shaped to provide adequate clot-trapping area while permitting sufficient blood flow. Also known are filters formed of various loops of wire, including some designed to partially deform the vessel wall in which they are implanted.

Typically, endoluminal devices such as vena cava filters and stents expand by one mechanism or another, not by some combination. That is, balloon expandable devices are not also self-expanding, and self-expanding devices are not typically balloon expandable. While it is known to "model" self-expanding stents to conform the anatomy in which the stent has been deployed to a shape suitable for accepting the stent, such modeling does not vary the diameter of the stent, but rather varies the anatomy. For example, a self-expanding stent deployed in an artery having a plaque deposit on the walls may be modeled to crush the plaque deposit down.

Self-expanding devices typically are known for excellent crush resistance, potential for longitudinal flexibility, for exerting a constant outward force to maintain an open vessel, and for having the capacity to elastically expand and contract with the blood vessel. On the other hand, self-expanding devices typically must be sized accurately to provide acceptable outward force and cannot generally be manipulated to a larger diameter after deployment. Self-expanding devices are typically made of a superelastic material, such as a superelastic grade of nitinol, which has a limited x-ray visibility.

Balloon-expandable devices are typically capable of expansion into a range of sizes, and typically are relatively stiff and inflexible. Balloon-expandable devices do not

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typically recover if crushed in the body, however, and are rigid bodies that do not expand and contract with motion of the vessel. Balloon-expandable devices are also typically longitudinally rigid, limiting delivery through tortuous anatomy. Balloon-expandable devices are made of a plastically deformable metal such as stainless steel, platinum, or a plastically deformable grade of nitinol.

In light of the advantages and disadvantages of self-expanding and balloonexpandable technologies as currently known, there is a need in the art for technology bridging the advantages of both.

#### SUMMARY OF THE INVENTION

It is to be understood that both the foregoing general description and the following detailed description are exemplary, but are not restrictive, of the invention. The invention comprises an endoluminal device, such as a stent or a vena cava filter, comprising at least one superelastic section and at least one plastically deformable section. The device typically has a first constrained diameter, a second fully-self-expanded diameter, and a third fully-forcibly-expanded diameter, wherein the third diameter is greater than the second diameter and the second diameter is greater than the first diameter.

The device may comprise a plurality of filaments including one or more superelastic filaments and one or more plastically deformable filaments, particularly where one or more of the superelastic filaments extends longitudinally substantially parallel to one or more of the plastically deformable filaments along the length of the stent. In the alternative, each superelastic section and/or each plastically deformable section may comprise a sheet or a longitudinally severed tube cut by a laser or other precision cutting mechanism.

Each superelastic filament may be connected along one or more longitudinal portions thereof to another superelastic filament, another columnar unit of the same superelastic filament, one or more plastically deformable filaments, or a combination thereof. Similarly, each plastically deformable filament may be connected along one or more longitudinal portions thereof to another plastically deformable filament, another columnar unit of the same plastically deformable filament, one or more superelastic filaments, or a combination thereof. The longitudinal portions may be connected at a joint by brazing, welding, adhesive bonding, or suturing.

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The superelastic filament or filaments preferably comprise a superelastic grade of nitinol, such as thermal shape memory nitinol. The plastically deformable filament or filaments preferably comprise gold, platinum, tantalum, titanium, stainless steel, tungsten, a cobalt alloy, a nickel alloy, a titanium alloy, or a combination thereof, such as a plastically deformable grade of nitinol or a composite nitinol wire having a core of a plastically deformable material (hereinafter referred to as a "deformable core nitinol composite").

Each plastically deformable section may comprise a constrained portion of the superelastic section comprising a plastically deformable material, such as gold. For example, the constrained portion may comprise a combination of superelastic material and plastically deformable material, such as plastically deformable material plated onto superelastic material, a plastically deformable hypotube overlaid onto superelastic material, plastically deformable material ion implanted within superelastic material, or a composite of deformable material and superelastic material. In particular, the device may be cut from a composite comprising a sandwich having outer layers of nitinol and inner longitudinal or transverse stripes of plastically deformable material. In an alternative embodiment, the device may comprise one or more hoops in a zig-zag configuration of oppositely-pointing apex sections, where the plastically deformable sections comprise one or more apex sections comprising plastically deformable material. The plastically deformable apex sections on each of said hoops may be longitudinally aligned.

The device may also comprise a first tubular section comprising a superelastic section and a second tubular section comprising a plastically deformable section. Where the first and second tubular sections both comprise a combination of superelastic material and plastically deformable material, the plastically deformable section has a ratio of plastically deformable material to superelastic material that is greater than the ratio of plastically deformable material to superelastic material in the superelastic section. The device may comprise a plastically deformable section mounted between two superelastic sections each having different fully-self-expanded diameters, so that the device can be mounted in a tapered lumen and forcibly expanded to conform to the taper in the lumen in accordance with a method of this invention.

The invention also comprises a method of deploying an endoluminal device of this invention in a body lumen. The method comprises introducing the device into the body lumen with the device radially constrained in a first configuration having the first diameter; allowing the device to self-expand into a second configuration having a diameter less than or equal to the second diameter; and optionally forcibly expanding at least a portion of the device into a third configuration having a diameter greater than the second diameter and less than or equal to the fully-expanded diameter.

Where the device comprises a plastically-deformable tubular section and a superelastic tubular section, the method may comprise deploying the device in a lumen having a tapered portion. In such case, the method further comprises introducing the device with the device radially constrained in a first configuration in which each tubular section has a first diameter. Next, the device is allowed to expand such that the plastically deformable section aligns with the tapered portion of the lumen. Thus, the device self-expands into a second configuration in which the superelastic section has a second diameter greater than the first diameter and less than or equal to the fully-self-expanded diameters. The device is finally forcibly expanded into a third configuration in which the plastically deformable section has a variable diameter that conforms to the tapered section of the lumen.

#### BRIEF DESCRIPTION OF DRAWINGS

The invention is best understood from the following detailed description when read in connection with the accompanying drawing. It is emphasized that, according to common practice, the various features of the drawing are not to scale. On the contrary, the dimensions of the various features are arbitrarily expanded or reduced for clarity. Included in the drawing are the following figures:

- Fig. 1 is a side view illustration of an exemplary filamentary stent of the present invention cut longitudinally and laid flat, consisting of two filaments.
  - Fig. 2 is a side view illustration of an exemplary brazed joint.
- Fig. 3A is a side view illustration of an exemplary zig-zag stent hoop of the present invention cut longitudinally and laid flat, in a constrained configuration.
  - Fig. 3B is a side view illustration of the exemplary zig-zag stent hoop of Fig. 3A in a fully-self-expanded configuration.
  - Fig. 3C is a side view illustration of the exemplary zig-zag stent hoop of Fig. 3A in a fully-forcibly-expanded configuration.

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- Fig. 4A is a side view illustration of an exemplary filamentary stent of the present invention cut longitudinally and laid flat, comprising a plurality of superelastic and plastically deformable filaments, in a constrained configuration.
- Fig. 4B is a side view illustration of the exemplary filamentary stent of Fig. 4A in a fully-self-expanded configuration.
  - Fig. 4C is a side view illustration of the exemplary filamentary stent of Fig. 4A in a fully-forcibly-expanded configuration.
  - Fig. 4D is a top view illustration of the exemplary filamentary stent of Fig. 4A in the constrained configuration of Fig. 4A.
  - Fig. 4E is a top view illustration of the exemplary filamentary stent of Fig. 4A in the fully-self-expanded configuration of Fig. 4B.
  - Fig. 4F is a top view illustration of the exemplary filamentary stent of Fig. 4A in the fully-forcibly-expanded configuration of Fig. 4C.
  - Fig. 5A is a side view illustration of an exemplary laser-cut stent of the present invention cut longitudinally and laid flat, in a constrained configuration.
  - Fig. 5B is a side view illustration of the exemplary laser-cut stent of Fig. 5A in a fully-self-expanded configuration.
  - Fig. 5C is a side view illustration of the exemplary laser-cut stent of Fig. 5A in a fully-forcibly-expanded configuration.
- Fig. 6 is a side view illustration of an exemplary prosthesis cut longitudinally and laid flat, comprising a plurality of zig-zag stent hoops.
  - Fig. 7A is a perspective view of an exemplary composite sheet having continuous longitudinal stripes of plastically deformable material, used for making an exemplary stent embodiment of the present invention.
  - Fig. 7B is a perspective view of a tube formed from the composite sheet of Fig. 7A, after a rolling step.

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Fig. 7C is a perspective view of a composite tube comprising an inner and outer hypotube.

Fig. 7D is a perspective view of an exemplary composite sheet having transverse stripes of plastically deformable material, used for making an exemplary stent embodiment of the present invention.

Fig. 7E is a perspective view of a tube rolled from the composite sheet of Fig. 7D.

Fig. 7F is a plan view illustration of a pattern to be cut on the composite sheet of Fig. 7A.

Fig. 8A is a perspective view of an exemplary assembled composite tube and its individual components for making an exemplary stent embodiment of the present invention, prior to a tube drawing step.

Fig. 8B is a perspective view of a tube of Fig. 8A after the tube drawing step.

Fig. 9 is a longitudinal section illustration of an exemplary prosthesis deployed in a lumen.

Figs. 10 is a side view illustration of an exemplary prosthesis comprising plastically deformable elements only at opposite ends of the stent.

#### DETAILED DESCRIPTION OF INVENTION

The invention will next be illustrated with reference to the figures wherein similar numbers indicate the same elements in all figures. Such figures are intended to be illustrative rather than limiting and are included herewith to facilitate the explanation of the apparatus of the present invention.

Although the term "vena cava filter" technically refers to implantable filters placed in the vena cava -- the jugular or femoral veins, which are the typical locations for such filters -- the term as used herein refers to any type of implantable filter, regardless of the specific body lumen into which it is implanted. In particular, the anchoring assemblies or portions of vena cava filters shown in U.S. Patent Nos. 5,709,704 and 5,836,939 may be

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particularly well-suited for construction in accordance with this invention. Vena cava filters are deployed much in the same way as stents, and thus throughout this application the term "endoluminal device" is used to refer to both vena cava filters and stents. In many instances, however, examples refer specifically to stents, but should not be interpreted to be limited to stents alone. Thus, discussion of aspects of this invention with respect to stents should also be considered applicable to implantable filters and vice versa, where applicable.

Fig. 1 shows an exemplary stent embodiment according to the present invention wherein the tubular stent has been cut along a line parallel to the tubular stent axis and flattened. The architecture of stent 12 shown in Fig. 1 is substantially similar to the architectures described in U.S. Patent Nos. 5,354,308 and 5,395,390 to Simon *et al.*, which are incorporated herein by reference, and described further below. The structures described in the '308 and '390 Simon patents are merely examples of particular stent architectures, and are not intended to be limiting. Other U.S. patents and patent families claiming priority therefrom, incorporated herein by reference, that may be well-suited for use in this invention include:

5,019,090	Pinchuk
5,135,536	Hillstead
5,292,331	Boneau
5,282,824	Gianturco
5,354,308	Simon et al.
5,507,767	Maeda et al.
5,800,515	Nadal <i>et al</i> .

The above list of patents is only exemplary, not limiting, however, as any number of different stent architectures may be used to create stents in accordance with this invention.

Stent 12 comprises a plurality of filaments 15 and 17. Superelastic filaments 15 typically comprise a superelastic grade of nitinol, whereas plastically deformable filaments

17 typically comprise gold, platinum, tantalum, titanium, stainless steel, tungsten, a cobalt alloy, a nickel or titanium alloy, such as a plastically deformable grade of nitinol or deformable core nitinol composite, or a combination of any of the above. The filaments may in the alternative comprise bioabsorbable or biostable polymers, such as are known in the art.

As used herein, a "superelastic" material is one that may be deformed into a certain configuration without the material permanently taking on the deformed shape. For example, a thermal shape memory material may be deformed into a number of configurations, but will return to its memory shape upon temperature activation. A "plastically deformable" material, on the other hand, is a material that once deformed into a certain shape, keeps that shape indefinitely, until deformed again by some other force.

As described in the background section, stents are typically inserted into a body lumen from a remote location through an insertion point in the body through which an "introducer", containing the stent in a radially compressed configuration, is threaded and navigated through the body lumen to the deployment location, where the stent is deployed in a radially expanded configuration. As referred to herein, "distal" refers to the direction further away from the insertion point and "proximal" refers to the direction closer to the insertion point.

As shown in Fig. 1, stent 12 comprises a filamentary stent architecture having a plurality of polygonal, hexagonal cells 30 having as sides thereof straight axially-extending portions 32 joined together along a set of parallel lengths 34, the parallel lengths deviating from one another proximally and distally of each parallel length into diagonal lengths 36, such that each set of parallel lengths and attached diagonal lengths form a Y-shaped intersection 38. The polygonal cell stent architecture in stent 12 terminates at a set of parallel lengths 34a, which may be joined together by welding, brazing, or any means known in the art, and additionally may comprise a single filament merely bent 180 degrees back upon itself, as shown in Fig. 1. In the alternative, the ends may be wound into an end winding having a different stent architecture, such as is described in U.S. Patent Application Serial No. 09/442,165, filed November 16, 1999, assigned to the common assignee of the present invention, and incorporated herein by reference.

As shown in Fig. 1, stent 12 comprises two sections 14 and 16. Section 14 is a superelastic section and section 16 is a plastically deformable section. Each section 14 and

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16 extends longitudinally along length L of the stent. As shown in Fig. 1, stent 12 consists of two filaments: superelastic filament 15 and plastically deformable filament 17. For ease of illustration in Fig. 1, stent 12 is shown after having been slit longitudinally and flattened, cutting filament 15 into two sections. When stent 12 is in its normal tubular form, however, filament ends 15a and 15b are connected to one another as part of continuous filament 15. For ease of identification, plastically deformable filament 17 is shown in Fig. 1 shaded in gray. Throughout the figures, gray shading is used to distinguish the plastically deformable elements from superelastic elements.

Stent 12 has opposite ends 18 and 20. Each filament 15 and 17 longitudinally traverses length L of stent 12 from the end 18 to the end 20 in a plurality of columnar units 22. As used herein, the term "columnar unit" refers to a single pass of the filament from one of ends 18 or 20 to the opposite end. Thus, as shown in Fig. 1, filament 15 comprises ten columnar units and filament 17 comprises two columnar units. In an alternative embodiment, each columnar unit, each pair of columnar units, or any number of columnar units may comprise a separate filament. Where each columnar unit is a separate filament, as shown in Fig. 4C, each superelastic filament 15 and each plastically deformable filament 17 extends only once between end 18 and end 20. Thus, stent 40 as shown in Fig. 4C comprises four superelastic filaments and four plastically deformable filaments. In such a case, ends 15d and 15e, for example, may be welded, brazed, twisted, glued, sutured, or otherwise connected together by means other than by a 180-degree turn 15c of a continuous wire, such as shown in Fig. 1.

As shown in Fig. 1, superelastic filament 17 extends longitudinally substantially axially parallel to plastically deformable filament 15 along length L of stent 12. As used herein, "substantially parallel" is used to denote that despite the serpentine bending of each filament about axis *I*, the axes of each columnar unit are essentially parallel to one another when the stent is in an unconstrained configuration. "Essentially parallel" is used to acknowledge that some minimal deviation from parallel may be possible.

Superelastic filament 15 is connected along one or more longitudinal portions thereof (each pair of parallel lengths 34) to either another columnar unit of the same superelastic filament, such as at 34a or 34b, or to plastically deformable filament 17, such as at 34c. Similarly, plastically deformable filament 17 is connected along one or more longitudinal portions thereof to another columnar unit of the same plastically deformable

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filament, such as at 34d, or to superelastic filament 15, such as at 34c, or some combination thereof. Where the stent comprises multiple superelastic filaments 15 and multiple plastically deformable filaments 17, such as illustrated in Fig. 4C, each plastically deformable filament may be connected along one or more longitudinal portions thereof to another plastically deformable filament, such as at 34e, or to a superelastic filament, such as at 34f, and each superelastic filament may be connected along one or more longitudinal portions thereof to another superelastic filament, such as at 34g, or to a plastically deformable filament, such as at 34f.

The longitudinal portions of the filaments at each set of parallel lengths 34 may be connected to make a joint by any means known in the art, such as brazing, welding, adhesively bonding, or suturing. Welding, suturing, adhesive bonding, and brazing processes are well-known in the art. Where the joint is a welded joint, it is preferable that both the superelastic material and the plastically deformable material are grades of nitinol. A superelastic grade of nitinol is any grade that is in its Austenitic phase at the use temperature (essentially the temperature of the human body). A plastically deformable grade of nitinol is any grade that is in its Martenisic phase at the use temperature, such as Alloy M, manufactured by Shape Memory Application Inc., of San Jose, CA, having a typical Active Austenite Finish A<sub>f</sub> in the range of 45-95°C, or Alloys B (A<sub>f</sub> 45-80°C) or BH (A<sub>f</sub> 80-120°C), manufactured by Memry Corporation, of Brookfield, CT. Deformable core nitinol composites are also plastically deformable nitinol materials. Where the joint is a brazed joint, the preferred superelastic material is nitinol and the preferred plastically deformable material is stainless steel, a titanium alloy, or platinum.

A brazed joint may comprise any wire wrapped around the joint to provide mechanical support. An exemplary brazing method is described, below. As shown in Fig. 2, joint 21 comprises filaments 22 and 24 wrapped by coil 26. One of filaments 22 and 24 may be superelastic, and the other plastically deformable, or both filaments may be the same type of material. To form such a brazed joint, wires 22 and 24 are brought together so that they abut, and a brazing material, such as any standard brazing material known in the art, is melted to wet at least the portions of both wires that contact one another, and cooled to solidify. Coil 26 is then wrapped around the brazed, abutted filaments. Coil 26 may comprise a deformable metal or a thermal shape memory material, such as nitinol. Where a shape memory material is used, it may be first wound into a tight helical coil around a small diameter form and annealed to set its Austenite shape. It is then cooled to its Martensite

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configuration and unraveled to be re-coiled about the brazed, abutted filaments. A heating step then shrinks joining coil 26 back to its tight Austenite shape. A low-temperature brazing material may then be melted and solidified over the coil to fix the coil in its Austenite shape.

In the alternative, instead of a filamentary coil, a split hypotube may be annealed in a closed shape, cooled to its Martensite condition and opened to allow it to be slipped over abutted filaments 22 and 24, and heat shrunk back to its closed shape. The low temperature brazing material may then be used to seal the hypotube in the closed position.

In an alternative embodiment, shown in Figs. 3A-C, stent 30 may comprise a zig-zag stent architecture. As shown in Fig. 3C, a zig-zag stent architecture comprises a plurality of struts 300 joined at apex sections 302, with a plurality of alternating oppositely-pointed apex sections arranged into cylindrical hoops, shown in Figs. 3A-C after being longitudinally cut and flattened. Apex sections 302 pointing in a first direction can be referred to peaks 306 and apex sections pointing in the opposite direction can be referred to as valleys 308, with any circumferentially adjacent, oppositely pointing apex sections together with three adjacent struts 300 forming a zig-zag 310. The zig-zag architecture shown in Fig. 3 is merely one exemplary zig-zag embodiment, but is not intended to limit the invention thereto.

As shown in Fig. 3C, each plastically deformable section 16 comprises a constrained portion of superelastic section 14. Plastically deformable section 16 in Fig. 3C comprises a combination of plastically deformable material 320 and superelastic material (filament 15). Fig. 3C shows a single hoop in a zig-zag configuration, split and laid flat, of oppositely-pointing apex sections (peaks 306 and valleys 308), where each plastically deformable section 16 comprises a peak 306 comprising plastically deformable material. As shown in Fig. 3C, the combination comprises plastically deformable material overlying the superelastic material, such as made by plating deformable material, such as gold, onto the superelastic material, or by overlying a plastically deformable hypotube onto the superelastic material. The combination may instead comprise plastically deformable material ion implanted into superelastic material, or a composite of deformable material and superelastic material. Ion implantation processes are well-known in the art. A composite combination is described herein below.

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As shown in Fig. 6, a prosthesis 60 (shown split and laid flat) may comprise a plurality of stent hoops 61 attached to a graft 62, wherein peaks 306 on each hoop are longitudinally aligned with the peaks of the other hoops. The plastically deformable apex sections 16 on each of the hoops may then be longitudinally aligned with the plastically deformable apex sections on the other hoops.

Although shown as a filamentary stent in Figs. 1-4F, stent 12 may also comprise, at least in part, a hypotube or sheet cut by precision cutting means such as a laser or chemical etching, as shown in Figs. 5A-5C. While laser cutting is the most common such method for performing such a cutting step, as is known in the art, the structure is not limited to any particular cutting mechanism. As shown in Fig. 5C, superelastic section 14 of stent 50 comprises a laser-cut sheet or a longitudinally severed laser cut tube. The opposite edges 51 and 52 of superelastic section 14 are joined together by plastically deformable section 16. Plastically deformable section 16 may be a portion of a laser-cut sheet or longitudinally severed laser cut tube itself, in which each joint 21 between sections 14 and 16 may be attached by brazing, welding, adhesive bonding, or any method known in the art. Section 16 may instead be filamentary, such as for example, a filament having a zig-zag architecture stent (not shown) wound helically between edge-most cells 30a and edge-most cells 30b on adjacent opposite edges 51 and 52 of superelastic section 14. For example, the filament may be wound in order from cell 30a<sub>1</sub> to 30b<sub>2</sub> to 30a<sub>2</sub> to 30b<sub>3</sub> to 30a<sub>3</sub> to 30b<sub>3</sub> to 30a<sub>4</sub> and optionally back again to cell 30a<sub>1</sub> through all of the same cells in reverse order.

As shown in Fig. 5C, plastically deformable section 16 comprises at least one columnar unit 22 having a longitudinal zig-zag configuration disposed between opposite longitudinal edges 51 and 52 of single superelastic section 14. Where there are multiple superelastic sections 14, each plastically deformable section may be disposed between two superelastic sections, and each superelastic section may be disposed between two plastically deformable sections.

In an alternative embodiment, stent 50 may comprise a single composite hypotube having plastically-deformable longitudinal stripes that is cut to form a seamless stent. Figs. 7A-8B illustrate various methods for forming a composite hypotube. As shown in Fig. 7A, a composite sandwich sheet 750 may comprise a lower layer 752 and upper layer 754 comprising superelastic material, such as nitinol, and a plurality of continuous longitudinal stripes 756a-d comprising plastically deformable material. Plastically

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deformable stripes 756a-d may comprise high density metal such as Pt, Ta, Au, Pd, W and the like, which provide improved visibility under x-ray imaging. Stiffer metals, such as W, may be preferred to add more stiffness to the composite. Each stripe 756a-d may comprise the same material, different materials, or alternating stripes of two or more different materials may be used, such that, for example, stripes 756a and 756c may comprise a first material, while stripes 756b and 756d comprise another. The thickness of individual stripes 756a-d, spaces 757a-c between the stripes, and overall number of stripes, in conjunction with the materials of the stripes, all may be varied as desired to obtain desired composite characteristics. Stripes may be longitudinal as shown in Figs. 7A-7C, or transverse as shown in Figs. 7D-7E, and may be continuous or broken. Thus, for example, stripes 756a-d may extend from end to end of lower sheet 752, or may only be present at the ends and not in the middle, or in selected longitudinal sections to provide areas of balloon-expandability only where desired. Regardless of whether the stripes are continuous or broken, or longitudinal or transverse, all of the embodiments described herein can be described as comprising a layer of plastically deformable material in a non-continuous distribution sandwiched between layers of superelastic materials.

Composite sandwich sheet 750 may be fabricated by depositing stripes 756a-d on lower layer 752 or upper layer 754, the two layers placed together with the stripes in between, and then the sandwich rolled to a final thickness. The rolled sandwich may then be formed into a tube 751 with the ends joined together at weld 758, as shown in Fig. 7B. Layers 752 and 754 may be fused together with stripes 756a-d in between using a hot isostatic press prior to rolling. Stripes 756a-d may be deposited by electroplating, ion beam deposition, plasma spray, laser-assisted deposition, or any method known in the art. In particular, electro-polishing the surface of nitinol onto which the stripes are to be deposited (to remove oxides from the metal surface) has been found to be desirable prior to the deposition step. For gold electroplating, the use of an acid-based gold strike process is preferred, such as is described in U.S. Patent Serial No. 09/227,407 by Steven Taskovics et al., filed on January 8, 1999, assigned to the common assignee of this invention and incorporated herein by reference.

In another embodiment, shown in Fig. 7C, longitudinal stripes 756a-d may be electroplated onto an inner hypotube 760 and the striped inner hypotube placed inside an outer hypotube 762. The tube may then be drawn, as is known in the art, to create a continuous hypotube similar to that shown in Fig. 7B (absent weld 758). The non-continuous

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intermediate layer formed by stripes 756a-d allows for areas of nitinol to nitinol contact between the stripes to provide support during the drawing process, which is an advantage over the use of a continuous intermediate layer.

Although shown in Figs. 7A-7C with stripes 756a-d that run longitudinally, the stripes may also run transversely, as shown in Figs. 7D and 7E. Transverse stripes 756e and 756f may be formed between lower and upper sheets 752 and 754 respectively using the sheet fabrication process as shown in Fig. 7D and described above with respect to Figs. 7A-7B, or may be formed using the hypotube process as described above with respect to Fig. 7C. The final tube structure 764 may thus comprise a seamless tube, as shown in Fig. 7E, if manufactured using hypotubes and drawing the assembled hypotubes, or may have a longitudinal weld similar to weld 758 shown in Fig. 7B, if created as a rolled sheet and then joined at the ends. In either case, tube 764 comprises one or more longitudinal sections 766a and 766b comprising an inner ring of plastically deformable material. As with the longitudinal stripes, the transverse stripes may be continuous or broken, such as stripe 756f, which is broken into four pieces, 756f<sub>i-iv</sub> which results in a broken ring 766b of plastically deformable material. Thus, as shown in Fig. 7E, the resulting ring 766b essentially comprises a set of longitudinal stripes or plastically deformable material present only on end section 768 of tube 764.

Yet another process for forming a composite hypotube is illustrated with respect to Figs. 8A and 8B, using technology similar to technology used in the formation of superconducting wires. A relatively large diameter superelastic hypotube 802 and a relatively smaller diameter superelastic hypotube 804 are positioned concentrically and a number of even smaller diameter superelastic tubes 806 are packed between tubes 802 and 804, as shown in Fig. 8A. Into the core 807 of each tube 806 is placed a plastically deformable wire 808. Heat and pressure, such as in a hot isostatic press, may be used to consolidate the assembly 810 into a bonded structure. Then assembly 810 is drawn to a desired diameter, leaving the final tube structure 812 shown in Fig. 8B having longitudinal cylindrical portions of plastically deformable material embedded therein. A sacrificial center core (not shown) may be used within the center 805 of tube 804 to assist in drawing tube 812. The sacrificial center core may then be removed at a later time by etching or a "pull to shrink" process as is known in the art. Wires 808 may also be used in place of stripes 756 or 756a-d when forming the composite using the processes described with respect to Figs. 7A-7E. Although shown in Fig. 8A as having a hexagonal cross-section, tubes 806 may have

any geometric cross-section known in the art. In yet another embodiment, the tube structure 812 shown in Fig. 8B may be created by starting with a thick-walled hypotube (not shown), drilling longitudinal holes in the hypotube and filling them with wires 808, and then drawing the assembled composite to create structure 812.

It should be noted that although described herein with respect to the use of superelastic and plastically deformable composite materials, the above composite processes may also be used for forming composite materials using only plastically deformable materials. For example, instead of lower and upper sheets 752 and 754 or inner and outer hypotubes 760 and 762, respectively, comprising a superelastic material such as nitinol, these components may comprise stainless steel, a cobalt alloy, or the like. The provision of stripes of other materials may thus be used to provide improved radiopacity, increased or decreased stiffness, or some other desired material property. Furthermore, for composites of plastically deformable and superelastic materials, the choice of plastically deformable material and the amount thereof in the composite can be chosen to create a hybrid device that is both plastically deformable and superelastic as described herein, or can be chosen merely to enhance radiopacity without significantly affecting the superelastic properties of the superelastic material.

Referring now to Fig. 7F, to form the stent or vena cava filter, the resulting composite sandwich 750 (after rolling) may then be cut by etching or cutting as is known in the art, in a pattern 784 corresponding to the desired architecture of the device, such as the zig-zag architecture shown in Fig. 7F. The pattern may preferably be aligned so that the features of the architecture desired to be plastically deformable, such as every other upwardly-pointing apex 786, as shown in Fig. 7F, are aligned with plastically deformable stripes 756a-d within composite sheet 750. After cutting, the resulting flat pattern 784 may then be rolled into a tubular shape and ends 788a, 788b, and 788c welded together as 788a-788a, 788b-788b, and 788c-788c to form the desired device. In the alternative, the tubular composite shown in Figs. 7B or 7E may be cut in their tubular form to create the device. Although shown in Fig. 7F with the stripes oriented longitudinally with respect to the cylindrical orientation of the device, the stripes may also be transverse as shown in Figs. 7D and 7E.

Thus, in the embodiments using a composite sheet or tube, the plastically deformable section comprises a combination of plastically deformable and superelastic

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material, similar to the embodiments shown in Figs. 3A-C. In each embodiment, the critical high strain areas can comprise plastically deformable or superelastic sections. If all the high strain areas are superelastic, the device will be fully self-expanding. If all the high strain areas are plastically deformable, the device will be balloon expandable only. According to the present invention, some high strain areas comprise superelastic sections and some comprise plastically deformable areas such that the device is self-expanding with the ability to be fined tuned by balloon expansion.

In each of the embodiments shown herein, but referring now to Figs. 4D - 4F, the device has a first constrained diameter  $D_1$  for introduction into a body lumen, a second fully-self-expanded diameter  $D_2$  after initial deployment in a body lumen, and a third fully-forcibly-expanded diameter  $D_3$  after full forced expansion, such as with a balloon.  $D_3$  is greater than  $D_2$ , which is greater than  $D_1$ .

To deploy an endoluminal device of this invention in a body lumen, the method comprises first introducing the device into the body lumen with the device radially constrained in a first configuration, as shown in Figs. 3A, 4A, 4D, and 5A. As shown in Fig. 4D, the device in this configuration has a first diameter  $D_1$ . This radially constrained configuration typically arises after a constraining force F, is applied to the stent to collapse the device. The device in its constrained configuration is then typically loaded within an introducer for introduction into the body of a recipient.

After the introducer reaches the desired deployment position, the device is released from the sheath or other constraining means within the introducer, allowing the device to self-expand into a second configuration approaching or equal to the fully-self-expanded configuration shown in Figs 3B, 4B, and 5B. As shown in Fig. 4E, this fully-self-expanded configuration has a second diameter  $D_2$  greater than the first diameter  $D_1$ .

Then, optionally, a balloon is inserted within the circumference of the device and inflated to forcibly expand at least longitudinal portions of the device to a third configuration. As further described below, this third configuration may range between greater than the second configuration and less than or equal to the fully-forcibly-expanded configuration shown in Figs. 3C, 4C, and 5C.

As shown in Fig. 4F, the fully-forcibly-expanded configuration has a third diameter D<sub>3</sub> greater than the second diameter D<sub>2</sub>. During the forcible expansion step, the

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balloon may forcibly expand the device into a fourth, overexpanded configuration having a diameter greater than the third diameter, as the self-expanding section of the stent may become expanded past its resting state. After the balloon is deflated, however, the device will relax back to the third configuration.

It should be noted that in actual deployment of a device in the body, that the deployment method may be terminated after the device expands to the second configuration if the diameter in that configuration is deemed adequate to perform the desired function. In other instances, it may be necessary to fully expand the device into the third configuration. In most instances, however, the desired configuration may be intermediate the second and third configurations. Likewise, it should also be noted that the body lumen may partially constrain all or part of the device from fully expanding even to the fully-self-expanded configuration. Thus, the second configuration may comprise an expanded configuration wherein the device or portions of the device have a diameter somewhat less than  $D_2$ . Thus, the device in its finally deployed configuration may have a single diameter throughout, where that diameter ranges from somewhat less than  $D_2$  to less than or equal to  $D_3$ , or some longitudinal sections may have a diameter as small as somewhat less than  $D_2$ , other sections a diameter as great as  $D_3$ , and still other sections a diameter intermediate  $D_2$  and  $D_3$ .

The advantage of the present invention is that it can in fact be tailored to conform to the anatomy of the lumen in which it is deployed by deforming the plastically deformable section of the device without changing the characteristics of the superelastic section of the device. Thus, the invention comprises a self-expanding device that can be "fine tuned" by deforming the plastically deformable section to achieve optimum sizing. This may be particularly useful for adapting a device to a lumen having a non-round, more oval cross-section, an application for which self-expanding devices generally are considered deficient because of their tendency to deploy with a round cross-section. Additionally, to the extent that the plastically deformable materials can be selected having better x-ray visibility than superelastic materials, devices of the present invention may also have increased x-ray visibility without the need for adding special radiopaque markers.

Devices of this invention may be manufactured by any of the typical means known in the art, consistent with this invention. For example, filamentary devices may be wound upon a mandrel or formed into a sheet, then rolled into a cylindrical shape.

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Similarly, cut tube devices may be cut in sheet form and then rolled, or formed as a tube and cut in their tubular configuration.

Although described above with respect to devices wherein the superelastic section and the plastically deformable section extend longitudinally along the length of the device, each section may instead comprise tubular portions axially attached to one another, as Fig. 9 depicts device 700 deployed within a lumen 702 having a small shown in Fig. 9. diameter section 704, a tapered section 706, and a large diameter section 708. A first superelastic section 14a having a first self-expanded diameter  $d_1$  is deployed in section 704 and a second superelastic section 14b having a second self-expanded diameter  $d_2$  is deployed in section 708. It should be understood that the deployed self-expanded diameters  $d_1$  and  $d_2$ as constrained by lumen 702 may be slightly less than the fully self-expanded diameters (not shown) of section 14a and 14b. It should also be understood that portions 704 and 708 of lumen 702 may also be tapered somewhat, and that the self-expanded diameters may be variable from a more-lumen-constrained diameter at one end to a less-lumen-constrained diameter at the opposite end. Plastically deformable section 16a is deployed within tapered section 706. The presence of plastically deformable section 16a allows the taper of the device to be tailored specifically to the tapered diameter of tapered section 706 of lumen 702 such that the diameter of section 16a is essentially  $d_1$  at first end 720 and essentially  $d_2$  at second end 722. As used herein "essentially" means that the diameter at each end may be slightly greater or less than the respective diameter of the adjacent section. Of course, although shown in Fig. 9 with a plastically deformable section between two superelastic sections, the device may comprise two plastically deformable sections having a superelastic section therebetween, or any multiple of plastically deformable and superelastic sections, including merely one of each section.

Plastically deformable section 16 in device 700 may comprise all plastically deformable materials, or may be a combination of plastically deformable materials and superelastic materials as described above. For example, device 700 may comprise a zig-zag architecture as shown in Figs. 3A-3C and 6, comprising one or more filaments 15 wherein section 16a comprises plastically deformable material 320 that constrains at least portions of the filaments. Similarly, device 700 may comprise a cut tubular device in which section 16a comprises a composite portion of plastically deformable material and superelastic whereas sections 14a and 14b comprise superelastic material only, such as shown in Fig. 7E.

Depending on the difference in diameter between sections 704 and 708 of lumen 702,

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superelastic sections 14a and 14b may have the same fully-self-expanded diameter, or section 14b may have a larger fully-self-expanded diameter than section 14b. Thus, sections 14a, 14b, and 16 may comprise discrete sections attached together by any means known in the art, such as by suturing, welding, brazing, adhesively bonding, or attachment to a common graft. Sections 14a, 14b, and 16 may also be attached to one another by any of the means described in the aforementioned U.S. Patent Application Serial No. 09/442,165. In the alternative, device 700 may comprise a unitary superelastic construction in which sections 14a and 14b are identical in construction and fully-self-expanded diameter, and section 16 merely comprises constraining plastically deformable material combined with the superelastic material as described herein.

Finally, device 700 may comprise a device in which each of sections 14a, 14b, and 16a themselves comprise combinations of superelastic and plastically deformable subsections wherein each superelastic subsection and plastically deformable subsection extend longitudinally along the length of each section as shown, for example, in Fig. 1. Thus, section 16a may merely be relatively more plastically deformable than sections 14a and 14b. For example, as shown in Fig. 9, section 16a comprises more plastically deformable subsections than in sections 14a and 14b. Stated another way, section 16a has a higher ratio of plastically deformable material to superelastic material than in sections 14a and 14b.

Whereas device 700 as shown in Fig. 9 has a plastically deformable section in the middle, in some instances it may be preferable to have plastically deformable sections only on the ends. For example, as shown in Fig. 10, device 1000 has plastically deformable sections 1002 and 1003 on opposite ends and a superelastic section 1004 in the center. For simplicity, device 1000 is depicted as a solid cylinder in side view with each section clearly defined, but is understood to have any filamentary or cut tubular architecture known in the art, with each section not necessarily definable from external viewing. This configuration is particularly useful to address an existing problem with self-expanding stents that "jump off" the delivery catheters during deployment. The configuration of device 1000 assures that the ends stay in place on the balloon catheter until expanded to implant the device precisely where desired. As shown in Fig. 10, the device ends have longitudinal stripes 1006 of plastically deformable material, such as what would result on end 768 of tube structure 764 shown in Fig. 7E, for example, after a cutting or etching step to create the device architecture, as depicted in Fig. 7F. This allows the ends of the device to be tapered or flared as necessary to conform to the anatomy of the patient.

Although illustrated and described above with reference to certain specific embodiments, the present invention is nevertheless not intended to be limited to the details shown. Rather, various modifications may be made in the details within the scope and range of equivalents of the claims and without departing from the spirit of the invention.

#### What is claimed:

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- 1 1. An endoluminal device comprising at least one superelastic section and 2 at least one plastically deformable section.
- 2. The device of claim 1, wherein the plastically deformable section has a greater x-ray visibility than the superelastic section.
  - 3. The device of claim 1 having a length wherein each of the superelastic section and the plastically deformable section extend longitudinally along the length of the device.
    - 4. The device of claim 1 further comprising a plurality of filaments including one or more superelastic filaments and one or more plastically deformable filaments.
    - 5. The device of claim 4 having a length, wherein said one or more superelastic filaments extend longitudinally substantially parallel to said one or more plastically deformable filaments along the length of the stent.
    - 6. The device of claim 5 having a first end and a second end, wherein each of said superelastic filaments and said plastically deformable filaments extends only once from the first end to the second end of the stent.
    - 7. The device of claim 4 having a first end and a second end, wherein at least one of said superelastic filaments or deformable filaments longitudinally traverses the length of the stent from the first end to the second end in a plurality of columnar units.
    - 8. The device of claim 4 consisting of a single superelastic filament and a single plastically deformable filament.
  - 9. The device of claim 4, wherein each of said superelastic filaments is connected along one or more longitudinal portions thereof to another superelastic filament, another columnar unit of the same superelastic filament, one or more of said plastically deformable filaments, or a combination thereof, and each of said plastically deformable filaments is connected along one or more longitudinal portions thereof to another plastically

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- deformable filament, another columnar unit of the same plastically deformable filament, one or more of said superelastic filaments, or a combination thereof.
- 1 10. The device of claim 9, wherein the longitudinal portions are connected 2 at a joint by one of: a brazed connection, a weld, an adhesive bond, or a suture.
  - 11. The device of claim 9 further comprising one or more joints comprising: a first longitudinal portion of one of the superelastic filaments, a second longitudinal portion of one of the plastically deformable filaments abutting said first portion, and a joining coil wrapped about said first and second portions.
    - 12. The device of claim 11, wherein said superelastic filaments comprise a superelastic grade of nitinol; said plastically deformable filaments comprise a material selected from the group consisting of: gold, platinum, tantalum, titanium, stainless steel, tungsten, a nickel alloy, a cobalt alloy, a titanium alloy, and a combination thereof; and said brazed coil comprises a thermal shape memory grade of nitinol.
    - 13. The device of claim 1, wherein each said superelastic section comprises a precision-cut sheet or a longitudinally severed precision-cut tube.
    - 14. The device of claim 13, wherein each said plastically deformable section comprises at least one columnar unit having a zig-zag configuration disposed between two superelastic sections or between opposite longitudinal edges of a single superelastic section.
  - 15. The device of claim 14 consisting of a single plastically deformable section comprises a single columnar unit attached between opposite longitudinal edges of a single superelastic section.
  - 16. The device of claim 1, wherein each plastically deformable section comprises a combination of superelastic material and plastically deformable material wherein said plastically deformable material constrains the superelastic material.
  - 17. The device of claim 16, wherein said combination is selected from a group consisting of: plastically deformable material plated onto said superelastic material, a plastically deformable hypotube overlaid onto said superelastic material, ion implantation of

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- said plastically deformable material into said superelastic material, and a composite comprising said deformable material and said superelastic material.
  - 18. The device of claim 16, wherein the combination comprises a composite comprising plastically deformable material sandwiched between inner and outer layers of superelastic material.
- 1 19. The device of claim 16, wherein the plastically deformable material is 2 gold.
  - 20. The device of claim 16 further comprising one or more hoops in a zigzag configuration of oppositely-pointing apex sections, each plastically deformable section comprising one or more apex sections comprising said plastically deformable material.
  - 21. The device of claim 19 further comprising a plurality of hoops wherein the apex sections pointed in a first direction on each of said hoops are longitudinally aligned and the plastically deformable apex sections on each of said hoops are longitudinally aligned.
  - 22. The device of claim 1 having a first constrained diameter, a second fully-self-expanded diameter, and a third fully-forcibly-expanded diameter, wherein said third diameter is greater than said second diameter and said second diameter is greater than said first diameter.
  - 23. The device of claim 1, wherein each of said superelastic sections comprises nitinol and each of said plastically deformable sections comprises a plastically deformable material selected from the group consisting of: gold, platinum, tantalum, titanium, stainless steel, tungsten, palladium, a nickel alloy, a titanium alloy, a cobalt alloy, and a combination thereof.
  - 24. The device of claim 1, wherein the device is selected from the group consisting of: a stent and a vena cava filter.
- The device of claim 1, wherein said at least one superelastic section comprises a first tubular section and said at least one plastically deformable section comprises a second tubular section.

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- 26. The device of claim 25, wherein the first tubular section consists essentially of a superelastic material alone and the second tubular section consists essentially of plastically deformable material alone.
  - 27. The device of claim 25, wherein the second tubular section comprises a combination of superelastic material and plastically deformable material having a first ratio of plastically deformable material to superelastic material.
  - 28. The device of claim 27, wherein the device comprises two opposite end sections having a middle section therebetween, the middle section comprising the first tubular section, and the two opposite ends each comprising second tubular sections.
  - 29. The device of claim 28, wherein each end section comprises the plastically deformable material aligned in longitudinal stripes between stripes of superelastic material.
  - 30. The device of claim 27, wherein the first tubular section comprises a combination of superelastic material and plastically deformable material having a second ratio of plastically deformable material to superelastic material less than said first ratio.
  - 31. The device of claim 25 further comprising a third tubular section comprising a superelastic section, the second tubular section disposed longitudinally between the first tubular section and the third tubular section, the first tubular section having a first fully-self-expanded diameter and the second tubular section having a second fully-self-expanded diameter.
  - 32. The device of claim 31, wherein the first fully-self-expanded diameter is less than the second fully-self-expanded diameter, and the second tubular section has a fully-forcibly-expanded diameter at least as great as said second fully-self-expanded diameter.
  - 33. A method of manufacturing an endoluminal device having an architecture, said method comprising:
- (a) forming a composite comprising a first layer comprising a first material, a second layer comprising the first material, and an intermediate layer between the first and second layers comprising a second material in a non-continuous distribution; and

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cutting or etching away portions of the composite tube in a pattern to (b) 6 form the device architecture. 7 The method of claim 33, wherein step (a) comprises forming the 34. 1 composite as a sheet and rolling the sheet to a desired thickness. 2 The method of claim 34 further comprising forming the sheet into a 35. 1 tube prior to step (b). 2 36. The method of claim 34 further comprising forming the device 1 architecture into a tubular shape after step (b). 2 The method of claim 33, wherein step (a) comprises forming the 37. 1 composite as tube wherein the first layer is an inner annular layer and the second layer is an outer annular layer and the intermediate layer is an annular layer between the inner and outer layers. The method of claim 33 wherein the non-continuous distribution 38. comprises a continuous longitudinal stripe, a non-continuous longitudinal stripe, a continuous . 3 transverse stripe, or a non-continuous transverse rings. 1 A method of deploying an endoluminal device in a body lumen, the 39. 2 device comprising at least one superelastic section and at least one plastically deformable section, the method comprising: introducing the device into the body lumen with the device radially (a) 4 constrained in a first configuration having a first diameter; 5 allowing the device to self-expand into a second configuration having a (b) 6 second diameter greater than the first diameter and less than or equal to a fully-self-expanded 7 diameter; and optionally, 8 forcibly expanding the device into a third configuration in which at 9 (c)

least one longitudinal portion of said device has a third diameter greater than said second

diameter and equal to or less than a fully-forcibly-expanded diameter.

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1	40. The method of claim 39 wherein step (c) comprises using a balloon to
2	forcibly expand said device into said third configuration.
1 2 3 4	41. The method of claim 40 wherein step (c) further comprises using said balloon to forcibly expand at least portions of said device into a fourth, intermediate configuration having a fourth, overexpanded diameter greater than said fully-forcibly-expanded diameter, and then allowing said device to relax to said third configuration.
1	42. The method of claim 39 wherein the device comprises a first, tubular
1	section comprising one of the superelastic sections and a second tubular section comprising
2	one of the plastically-deformable sections, the first tubular section having a first fully-self-
4	expanded diameter and the second tubular section having a fully-forcibly expanded diameter
	greater than the first fully-self-expanded diameter, the method further comprising:
6	in step (a) introducing the device into the body lumen with the device radially
Ż	constrained in the first configuration in which each tubular section has the first diameter;
	in step (b) allowing the device to self-expand into the second configuration in
	which the first tubular section has the second diameter; and
2 2 11 15 12	
	in step (c) forcibly expanding the device into the third configuration in which
1 <b>1</b>	the second tubular section has a diameter greater than the second diameter of the first tubular
10 11 12	section.
1	43. The method of claim 42, wherein the device is deployed in a lumen
2	comprising a tapered portion, the method further comprising:
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3	in step (b) allowing the device to expand in a location wherein the second
4	tubular section is aligned with the tapered portion of the lumen; and
5	in step (c) forcibly expanding said second tubular section to conform to said
6	tapered portion of the lumen such that the second tubular section comprises a variable
7	diameter expanding from essentially the second diameter of the first tubular section at a first
8	end to larger diameter at a second end.
1	44. The method of claim 42, wherein the device has a middle section and

two opposite end sections, the first tubular section comprises the middle section, the end

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- sections each comprise second tubular sections, the device is introduced into the body on a balloon catheter, and in step (b) the second configuration comprises a configuration wherein the second tubular sections remain in contact with the balloon catheter.
  - 45. The method of claim 42, wherein the device has a middle section and two opposite end sections, the first tubular section comprises the middle section, the end sections each comprise second tubular sections, and the third configuration into which the second tubular section is forcibly expanded in step (c) comprises a configuration wherein one or both end sections are tapered.
  - 46. The method of claim 39 wherein the device comprises a first, tubular section comprising one of the superelastic sections, a second tubular section comprising one of the plastically-deformable sections, and a third tubular section comprising one of said superelastic sections, the second tubular section disposed longitudinally between the first tubular section and the third tubular section, the first tubular section having a first fully-self-expanded diameter, the third tubular section having a second fully-self-expanded diameter greater than or equal to the first fully-self-expanded diameter, and the second tubular section having a fully-forcibly expanded diameter at least as great as the second fully-self-expanded diameter, the method further comprising:

in step (a) introducing the device into the body lumen with the device radially constrained in the first configuration in which each tubular section has a first diameter;

in step (b) allowing the device to self-expand into the second configuration in which the first and third tubular sections each have respective second diameters greater than the respective first diameters and less than or equal to the respective fully-self-expanded diameters; and

in step (c) forcibly expanding the device into the third configuration in which the second tubular section has a diameter greater than said second diameter of the first tubular section.

47. The method of claim 46 wherein said third tubular section has a greater fully-self-expanded diameter than said first tubular section, and wherein the device is deployed in a lumen comprising a smaller diameter portion, a larger diameter portion greater

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- than said smaller diameter portion, and a tapered portion between said smaller diameter portion and said larger diameter portion, the method further comprising:
  - in step (b) allowing the device to expand in a location wherein the first tubular section is aligned with the smaller diameter portion of the lumen, the second tubular section is aligned with the tapered portion of the lumen, and the third tubular section is aligned with the larger diameter portion of the lumen; and

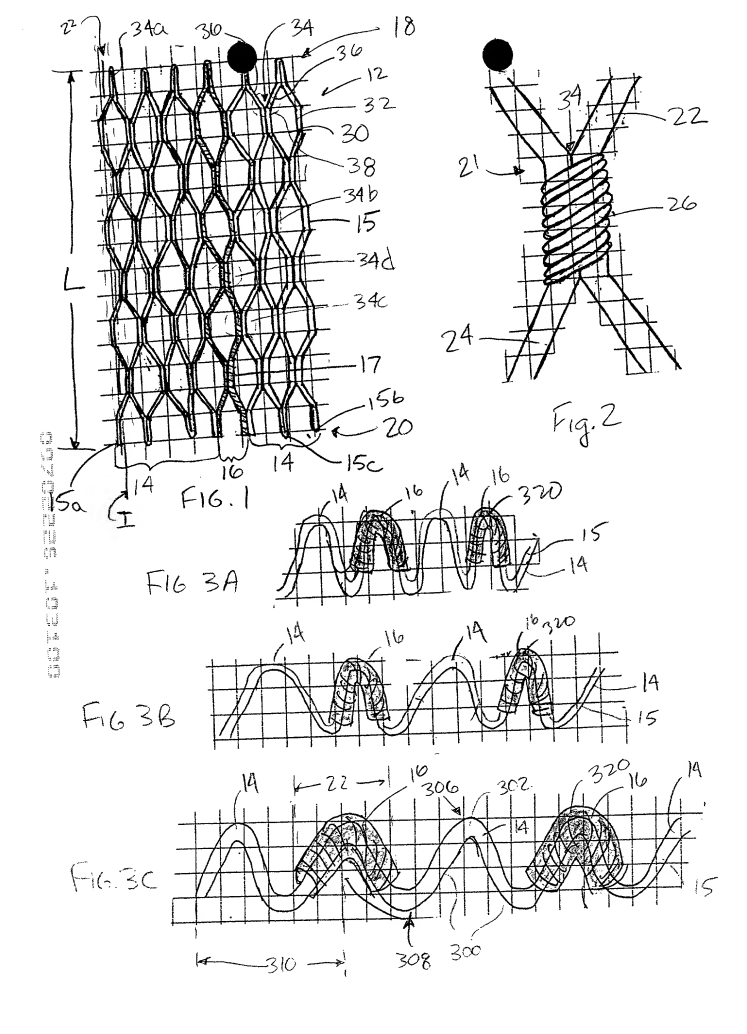
in step (c) forcibly expanding said second tubular section to conform to said tapered portion of the lumen such that the second tubular section comprises a variable diameter ranging from essentially the second diameter of the first tubular section at a first end to essentially the second diameter of the third tubular section at a second end.

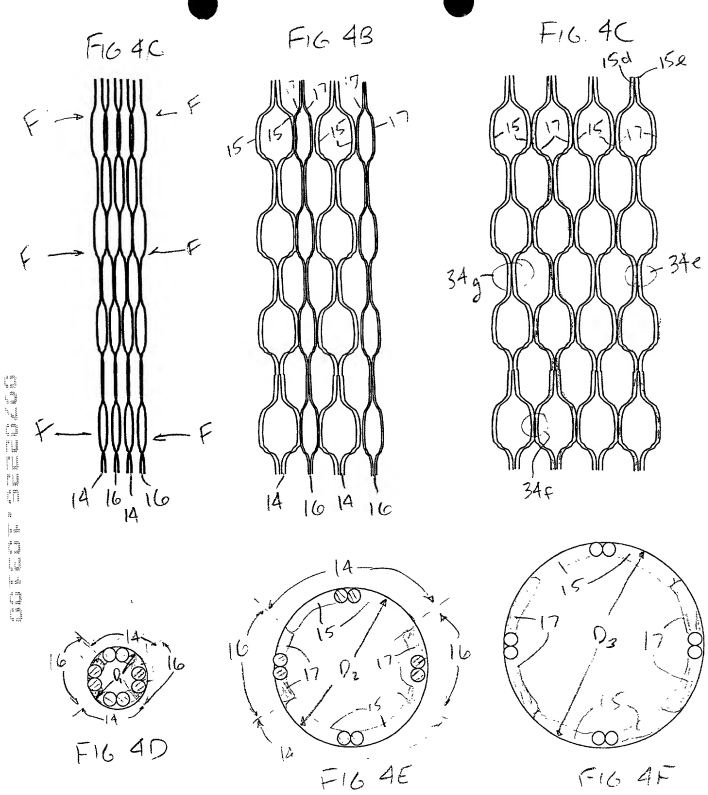
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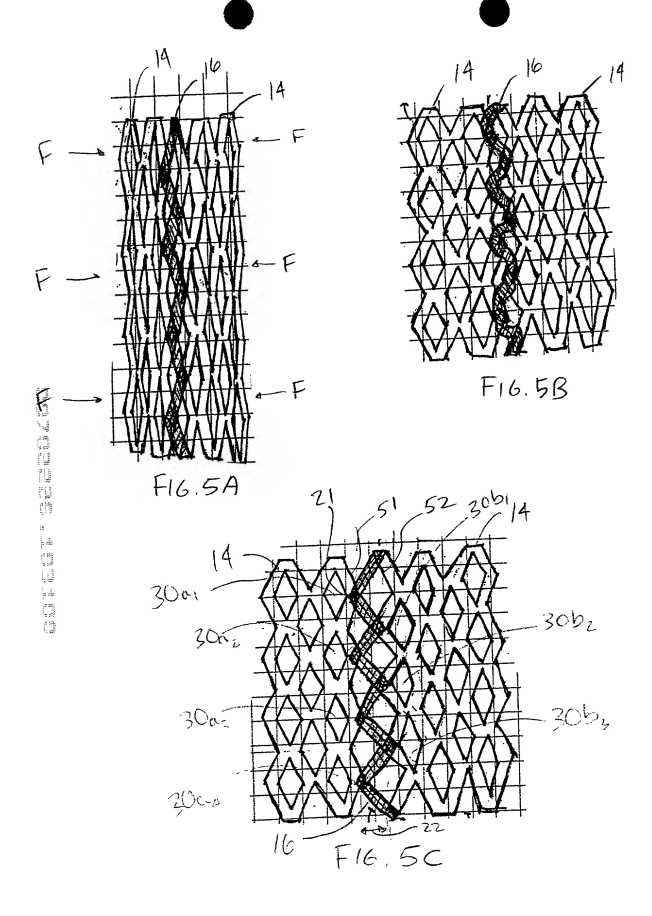
### COMBINATION SELF-EXPANDABLE, BALLOON-EXPANDABLE ENDOLUMINAL STENT

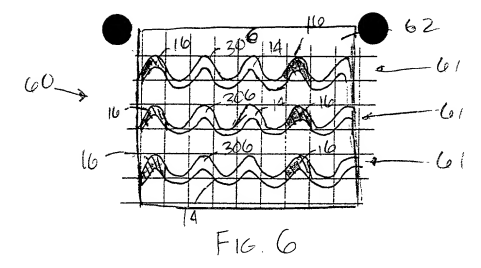
#### **ABSTRACT**

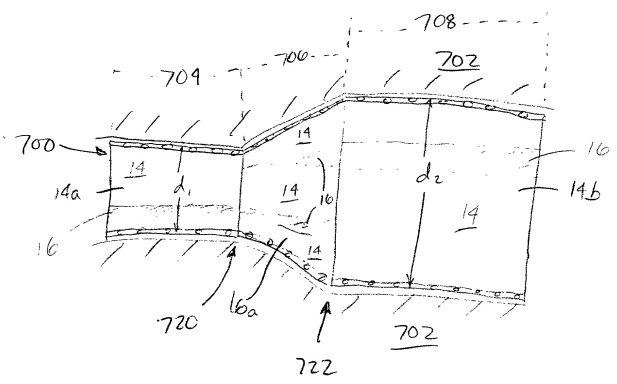
An endoluminal device, such as a stent or a vena cava filter, comprising at least one superelastic section and at least one plastically deformable section. The superelastic section may comprise, for example, a superelastic grade of nitinol, whereas the plastically deformable section may comprise, for example, gold, platinum, tantalum, titanium, stainless steel, tungsten, a nickel alloy, a cobalt alloy, a titanium alloy, or a combination thereof. Each plastically deformable section may merely comprise a constrained portion of the superelastic section comprising a plastically deformable material, such as gold. The device enables deployment by a method comprising introducing the device into a body lumen with the device radially constrained in a first configuration having a first diameter; allowing the device to self-expand into a second configuration having a second diameter less than or equal to a fully-self-expanded diameter; and then optionally "fine-tuning" the device by forcibly expanding the device into a third configuration having a third diameter in a range between the second diameter and less than or equal to a fully-forcibly-expanded diameter. The superelastic and plastically deformable sections may be tubular sections placed end-to-end, such that the plastically deformable section can be conformed to fit a tapered section of a lumen.











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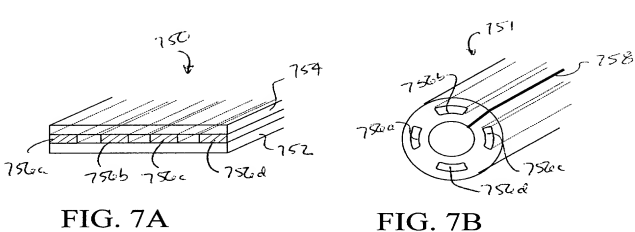
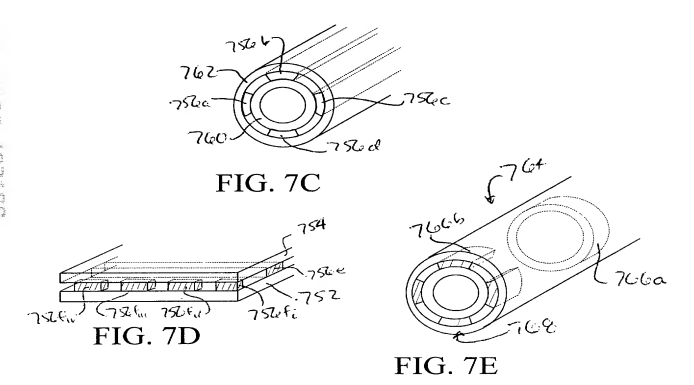
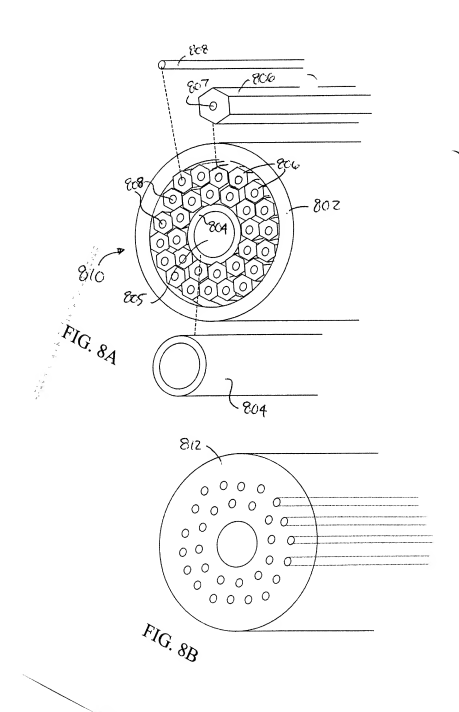


FIG. 7B





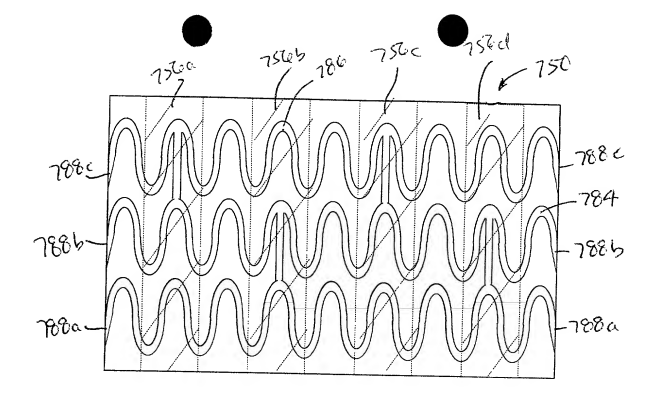


FIG.7F

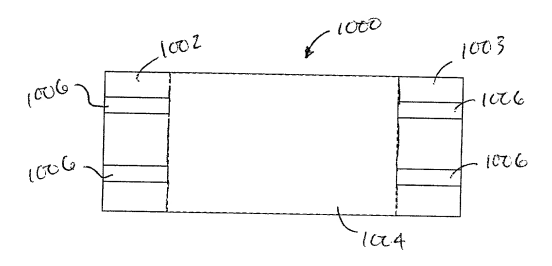


FIG. 10

# Declaration and Power of Attorney For Patent Application English Language Declaration

As a below named inventor, I hereby declare that:								
My residence, post office address and citizenship are as stated below next to my name,								
first and joint inventor and for which a pater COMBINATION SELI	r (if plural names are lis nt is sought on the inver F-EXPANDABLE, BALI	entor (if only one name is listed below) or sted below) of the subject matter which is ntion entitled LOON-EXPANDABLE ENDOLUMINAL unless the following box is checked:	s claimed					
and was amende I hereby state that I h	pplication Number or Po ed on (if applicat nave reviewed and unde	CT International Application Number ble). erstand the contents of the above identif nendment referred to above.						
I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR § 1.56.  I hereby claim foreign priority benefits under 35 U.S.C. §119(a)-(d) or § 365(b) of any foreign application(s) for patent or inventor's certificate, or § 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below by checking the box, any foreign application for patent or inventor's certificate, or PCT International application having a filing date before that of the application on which priority is claimed:  Prior Foreign Application(s)								
#								
(Number)	(Country)	(Day/Month/Year Filed)						
(Number)	(Country)	(Day/Month/Year Filed)						
I hereby claim the benefit under 35 U.S.C. § 119(e) of any United States provisional application(s) listed below.								
(Application Number)	(Filing Date)							
(Application Number)	(Filing Date)							
PCT International ap	oplication designating to the claims of this applications	§ 120 of any United States application( the United States, listed below and, ins cation is not disclosed in the prior Un	sofar as the subject					

international filing date of this application:

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09/362,261		July 28, 1999	Pending					
(Application Number)		(Filing Date)		nted, pending, abandoned	i)			
Í								
(Application Number)		(Filing Date)	(Status - pater	nted, pending, abandoned	d)			
agent(s) to prose	POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith:							
Paul F. Prestia Allan Ratner Andrew L. Ney Kenneth N. Nigon Kevin R. Casey Benjamin E. Leace James C. Simmons	Reg. No. 23,031 Reg. No. 19,717 Reg. No. 20,300 Reg. No. 31,549 Reg. No. 32,117 Reg. No. 33,412 Reg. No. 24,842	Lawrence E. Ashery Christopher R. Lewis Robert L. Andersen Joshua L. Cohen Daniel N. Calder Louis W. Beardell, Jr. Jacques L. Etkowicz	Reg. No. 34,515 Reg. No. 36,201 Reg. No. 25,771 Reg. No. 38,040 Reg. No. 27,424 Reg. No. 40,506 Reg. No. 41,738	Jack J. Jankovitz Jonathan H. Spadt Christopher I. Halliday Scott A. Mckeown	Reg. No. 42,690 Reg. No. 45,122 Reg. No. 42,621 Reg. No. 42,866			
Address all correspondence to: Paul F. Prestia Ratner & Prestia, Suite 301, One Westlakes, Berwyn, P.O. Box 980, Valley Forge, PA 19482-0980 Address all telephone calls to: Paul F. Prestia at (610) 407-0700.  I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.  Full name of sole or first inventor (given name, family name) Steven E. Walak  Inventor's signature  Date  10/26/00								
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Full name of second joint inventor, if any (given name, family name)								
Second Inventor's sig	nature			Date				
Residence								
Citizenship	· <del></del>							
Post Office Address	Post Office Address							
Additional inve	entors are being na	amed on separately number	red sheets attached	hereto.				